olefin-arene [2 + 2] cycloaddition¹⁵ followed by ring expansion and deprotonation. Thus, the nature of the photoaddition pathways followed by 1 appears to be critically dependent on the electron-donating ability of the olefin in a way strongly suggestive of electron-transfer mechanisms for pyrrolidine ether and olefin formation. Thus, competition between addition and cycloaddition might be regulated at an olefin-iminium salt exciplex stage by the relative magnitudes of k_{et} and k_{2+2} . Also, it is quite likely that the intramolecular version of this new photochemical reaction¹ observed for N-allyliminium salt systems also follows an electron-transfer mechanism, Further studies are underway to probe further the mechanistic details and synthetic potential of these reactions.

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Stereochemistry of Free-Radical Substitution on the Peroxide Bond

Sir:

The stereochemical course of the $S_H 2$ reaction has been the subject of intensive theoretical interest for some 40 years.¹⁻³ Calculations suggest^{1,3} that homolytic substitution in simple three-atom systems occurs via a colinear geometry and scattering experiments support the linear arrangement for deuterium-halogen exchange.⁴ Examples of experiments designed to provide stereochemical information about the S_H2 reaction in more complicated organic systems are rare and studies have been limited to halogen atom substitution on cyclopropane carbon for the first row elements.⁵⁻⁷ Homolytic substitution on phosphorus⁸ and sulfur⁹ has been studied and inversion of configuration is generally observed, although the possibility of a metastable radical-addition intermediate may complicate the interpretation of these results.

Carbon radical attack on the peroxide bond represents an important pathway in the autoxidation of olefins. For example, intramolecular carbon radical substitution (S_Hi) on the peroxide bond initiates the unzipping of styrene-oxygen copolymer.¹⁰ Because of the importance of this reaction in the oxi-



dation of polymers and natural products such as polyunsaturated lipids, we have initiated a study directed toward determining the stereochemical preference of carbon radical substitution on peroxide. We report here results of studies of the S_{H} i reaction in which the orientation of the attacking radical with respect to the peroxide bond is systematically varied. This approach gives information about the stereochemical preference of the substitution reaction and the results suggest that a "back-side attack" of the carbon radical on the peroxide bond is required.

The β -bromo peroxides 1-4 were prepared by reaction of



the corresponding mercuri bromides^{11,12} with molecular bromine.¹³ Compounds 1 and 2 are formed as three and erythro diastereomers and 4 consists of a mixture of cis and trans isomers as prepared,¹² These diastereomers can all be separated by high pressure liquid chromatography (HPLC) on μ -Porasil with 5% ethyl acetate-hexane.

The bromo peroxides 1-4 were reacted with 1.0 equiv of tributyltin hydride in benzene at 25 °C with 2-5% tert-butyl hyponitrite initiator present.¹⁴ For example, reaction of 1 (threo or erythro) with 0.455 M tin hydride leads to a mixture of cyclic peroxide 5 and epoxy alcohol 6 in a 5:6 ratio of 82:18.



The epoxy alcohol 6 was formed as a 3:1 mixture of the trans and cis geometric isomers and this product distribution was independent of the stereochemistry (threo or erythro) of the starting β -bromo peroxide. The results of analogous reactions of 2, 3, and 4 with tributyltin hydride led to mixtures of the corresponding cyclic peroxides and epoxy alcohols.¹⁵ The epoxy alcohol 7 derived from 2 rearranges under the conditions of analysis to the furan and pyran products 8 and 9.7 was prepared independently from the corresponding olefin and was converted into 8 and 9 by traces of acid.

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Table I. Product Distribution of 0.455 M Bromo Peroxide-Tin Hydride Reactions

bromo peroxide	peroxide, % ^a	epoxy Alcohol, % ^{<i>a,b</i>}
1	82	18
2	17	83
3	92	8
4	100	0

^a Product distributions are normalized to 100%; product accounted for was >80%. ^b Total epoxy alcohol 7 and furan and pyran 8 and 9 were analyzed.

Table II, r^a and k_{SHi} Values Found for Radicals 11-14

radical	r	k _{SHi} , s ⁻¹
11	0.09	7.5×10^{4}
12	1.05	8.7×10^{5}
13	0.01	1×10^{4}
14	<10-6	<1

$$a r = k_{\rm SHi}/k_{\rm H}$$

In Table I is presented the product composition for reaction of the bromo peroxides 1-4 with 0.455 M tributyltin hydride. Product accountability was high (85-100%) with reactions run with tin hydride concentrations >0.05 M.

The mechanism presented in Scheme I is consistent with the products observed.¹⁶ Further, for 1, 2, and 3 the product dis-

Scheme I

tribution varies as a function of Bu₃SnH concentration as is required by this mechanism. For example, the percent yield of peroxide 10 derived from 3 is 92, 82, 78, 66, and 55% as the tin hydride concentration is reduced from 0.455 M to 0.2, 0.1, 0.05, and 0.02 M. On the other hand, no epoxy alcohol could be observed in the reaction of 4 with Bu₃SnH with concentrations of hydride as low as 0.01 M.

With the mechanism presented in Scheme I as a format, a kinetic expression¹⁶⁻¹⁸ can be derived that gives a rate ratio of $r = k_{SHi}/k_H$ for the radicals 11–14 derived from 1–4. Further, since values of $k_{\rm H}$ are known,¹⁹ the $k_{\rm SHi}$ rate constants can be calculated (see Table II).

We suggest¹¹ that the dihedral angle ϕ must be 180°, or nearly so, for maximum S_Hi reactivity. For radical 12, derived from 2, a chair conformation²⁰ allows an equatorial radical to attack the peroxide bond from the back side. For the more planar dioxolanyl radicals 11 and 13 (from 1 and 3), it is difficult to adopt conformations with the 180° preferred angle for substitution, and the rates for substitution are thus one to two orders of magnitudes slower than $k_{\rm SHi}$ for 12. The endocyclic radical 14 formed from the seven-membered-ring peroxide, 4, is constrained to attack the peroxide bond from the side ($\phi < 70-100^\circ$) rather than via the back-side pathway and, as a consequence, no detectable S_{Hi} reaction is observed. It should be noted that the transition states for S_{Hi} reaction of the radicals 12 and 14 are isomeric but that the rate difference for substitution between these two radicals is $>10^6$ s⁻¹. This



observation supports the notion that the triangular transition state (side approach as in 14) is not favored¹⁻³ and points to a preferential colinear, or back-side, substitution process.

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- (21) NIH Research Career Development Awardee, 1977-1982. (b) NIH Pharmacology Trainee.

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Chemistry at Liquid-Liquid Interfaces. Evidence for an S_N1 Reaction Occurring at a Toluene-Water Interface¹

Sir:

Although many types of chemical and biochemical reactions are believed to take place at liquid-liquid interfaces, few examples have been well documented. It is not surprising, therefore, that only a limited understanding exists of how reactions at interfaces differ from comparable reactions carried